

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 163396

TO: Tamthom Truong

Location: REM/5C19/5C18

Art Unit: 1624

Wednesday, August 24, 2005

Case Serial Number: 10/083245

From: John DiNatale

Location: Biotech-Chem Library

REM-1B65

Phone: (571)272-2557

john.dinatale@uspto.gov

Search Notes

Examiner Truong,

See attached results.

If you have any questions about this search feel free to contact me at any time. Please note that in the broader search, the number of intervening carbon atoms (between the nitrogen atoms) was intentionally set to "1 to 6" in order to have a more conservative interpretation of your search request.

Thank you for using STIC search services!

John DiNatale Technical Information Specialist STIC Biotech/Chem Library (571)272-2557





STIC SEARCH RESULTS FEEDBACK FORM

Biotech-Chem Library

Questions about the scope or the results of the search? Contact the searcher or contact:

Mary Hale, Information Branch Supervisor Remsen Bldg. 01 D86 571-272-2507

Voluntary Results Feedback Form
> I am an examiner in Workgroup: Example: 1610
> Relevant prior art found, search results used as follows:
☐ 102 rejection
☐ 103 rejection
☐ Cited as being of interest.
☐ Helped examiner better understand the invention.
Helped examiner better understand the state of the art in their technology.
Types of relevant prior art found:
☐ Foreign Patent(s)
 Non-Patent Literature (journal articles, conference proceedings, new product announcements etc.)
> Relevant prior art not found:
Results verified the lack of relevant prior art (helped determine patentability).
Results were not useful in determining patentability or understanding the invention.
Comments:

Diep off or send completed forms to STIC-Blotsch-Chem Library Remsen Eldg.



> file registry FILE 'REGISTRY' ENTERED AT 14:32:51 ON 24 AUG 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 23 AUG 2005 HIGHEST RN 861509-89-9 DICTIONARY FILE UPDATES: 23 AUG 2005 HIGHEST RN 861509-89-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See ${\tt HELP\ SLIMITS}$ for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> file medline FILE 'MEDLINE' ENTERED AT 14:32:55 ON 24 AUG 2005

FILE LAST UPDATED: 23 AUG 2005 (20050823/UP). FILE COVERS 1950 TO DATE.

On December 19, 2004, the 2005 MeSH terms were loaded.

The MEDLINE reload for 2005 is now available. For details enter HELP RLOAD at an arrow promt (=>). See also:

http://www.nlm.nih.gov/mesh/ http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html

OLDMEDLINE now back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2005 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> file embase FILE 'EMBASE' ENTERED AT 14:33:00 ON 24 AUG 2005 COPYRIGHT (C) 2005 Elsevier Inc. All rights reserved.

FILE COVERS 1974 TO 18 Aug 2005 (20050818/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> file biosis

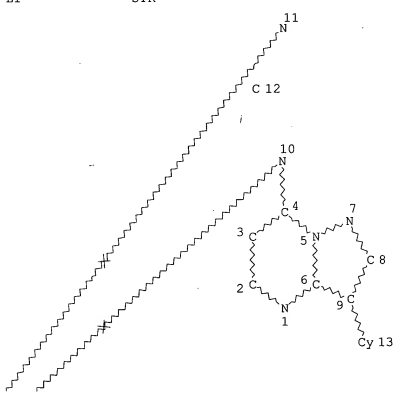
FILE 'BIOSIS' ENTERED AT 14:33:04 ON 24 AUG 2005 Copyright (c) 2005 The Thomson Corporation

FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 17 August 2005 (20050817/ED)

FILE RELOADED: 19 October 2003.

=> d stat que L14



Page 1-A

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Page 2-A
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GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

435 SEA FILE=REGISTRY SSS FUL L1

L14 0 SEA L6 => file beilstein FILE 'BEILSTEIN' ENTERED AT 14:33:37 ON 24 AUG 2005
COPYRIGHT (c) 2005 Beilstein-Institut zur Foerderung der Chemischen Wissenschaften licensed to Beilstein GmbH and MDL Information Systems GmbH

FILE RELOADED ON OCTOBER 20, 2002 FILE LAST UPDATED ON JUNE 29, 2005

FILE COVERS 1771 TO 2005.
*** FILE CONTAINS 9,271,550 SUBSTANCES ***

>>>PLEASE NOTE: Reaction Data and substance data are stored in separate documents and can not be searched together in one query. Reaction data for BEILSTEIN compounds may be displayed immediately with the display codes PRE (preparations) and REA (reactions). A substance answer set retrieved after the search for a chemical name, a compounds with available reaction information by combining with PRE/FA, REA/FA or more generally with RX/FA. The BEILSTEIN Registry Number (BRN) is the link between a BEILSTEIN compound and belonging reactions. For mo detailed reaction searches BRNs can be searched as reaction partner BRNs Reactant BRN (RX.RBRN) or Product BRN (RX.PBRN).<<<

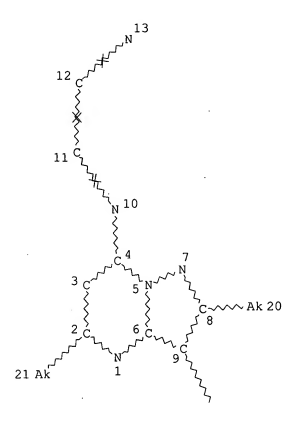
>>> FOR SEARCHING PREPARATIONS SEE HELP PRE <<<

- * PLEASE NOTE THAT THERE ARE NO FORMATS FREE OF COST.
- * SET NOTICE FEATURE: THE COST ESTIMATES CALCULATED FOR SET NOTICE
- * ARE BASED ON THE HIGHEST PRICE CATEGORY. THEREFORE; THESE
- * ESTIMATES MAY NOT REFLECT THE ACTUAL COSTS.
- * FOR PRICE INFORMATION SEE HELP COST

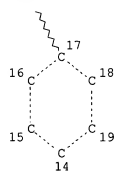
NEW

- * PATENT NUMBERS (PN) AND BABS ACCESSION NUMBERS (BABSAN) CAN NOW BE SEARCHED, SELECTED AND TRANSFERRED.
- * NEW DISPLAY FORMATS ALLREF, ALLP AND BABSAN SHOW ALL REFERENCES, ALL PATENT REFERENCES, OR ALL BABS ACCESSION NUMBERS FOR A COMPOUND AT A GLANCE.

=> d stat que L17 L8 STR



Page 1-A



Page 2-A NODE ATTRIBUTES: NSPEC IS R AT 1234567 IS R NSPEC AT NSPEC IS R ΑT NSPEC IS R ΑT NSPEC IS R AT NSPEC IS R AT NSPEC IS R ΑT 8 NSPEC IS R AT 9 **NSPEC** IS R AT ΑT 10 ·NSPEC IS RC ΑT 11 NSPEC IS RC AT 12 NSPEC IS RC **NSPEC** AT 13 IS RC

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                  AT
NSPEC
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                  ΑT
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NSPEC
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DEFAULT MLEVEL IS ATOM
MLEVEL IS CLASS AT 10 11 12 13 20 21
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M1-X6 C AT 20
ECOUNT IS M1-X6 C AT
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GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 21

STEREO ATTRIBUTES: NONE

L16 6 SEA FILE=BEILSTEIN SSS FUL L8

L17 5 SEA FILE=BEILSTEIN ABB=ON PLU=ON L16 NOT RN/FA

=> d qrd allref L17 1-5

L17 ANSWER 1 OF 5 BEILSTEIN COPYRIGHT 2005 BEILSTEIN MDL on STN

Beilstein Records (BRN): 1175562

Chemical Name (CN): N-(2,5-dimethyl-3-phenyl-pyrazolo<1,5-

a>pyrimidin-7-yl)-2-morpholin-4-yl-

acetamide

Autonom Name (AUN): N-(2,5-dimethyl-3-phenyl-pyrazolo<1,5-

a>pyrimidin-7-yl)-2-morpholin-4-yl-

acetamide

Molec. Formula (MF): C20 H23 N5 O2

Molecular Weight (MW): 365.43

Lawson Number (LN): 30824, 30310, 3379

Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 1117700 Tautomer ID (TAUTID): 1139153 Beilstein Citation (BSO): 5-27

Entry Date (DED): 1988/11/29
Update Date (DUPD): 1992/05/13

Field Availability:

Code	Name	Occurrence
======		
BRN	Beilstein Records	1
CN	Chemical Name	1
AUN	Autonomname	1
MF .	Molecular Formula	1
FW	Formular Weight	1
LN	Lawson Number	3
CTYPE	Compound Type	1
CONSID	Constitution ID	1
TAUTID	Tautomer ID	1
BSO	Beilstein Citation	1
DED	Entry Date	1
DUPD	Update Date	1
MP	Melting Point	1

This substance also occurs in Reaction Documents:

Code	Name	Occurrence
=======		=========
RX	Reaction Documents	1
RXPRO	Substance is Reaction Product	1

All References:

ALLREF

1. Patent: Shionogi and Co. Ltd. JP 7030335 1970, Chem. Abstr., 74(22872)

L17 ANSWER 2 OF 5 BEILSTEIN COPYRIGHT 2005 BEILSTEIN MDL on STN

Beilstein Records (BRN): 1164213

(2,5-dimethyl-3-phenyl-pyrazolo<1,5-Chemical Name (CN): a>pyrimidin-7-yl)-(2-morpholin-4-yl-ethyl)amine (2,5-dimethyl-3-phenyl-pyrazolo<1,5-Autonom Name (AUN): a>pyrimidin-7-yl)-(2-morpholin-4-yl-ethyl)amine C20 H25 N5 O Molec. Formula (MF): 351.45 Molecular Weight (MW): Lawson Number (LN): 30824, 30310, 3018 Compound Type (CTYPE): heterocyclic Constitution ID (CONSID): 1113261 Tautomer ID (TAUTID): 1135556 5-27 Beilstein Citation (BSO): 1988/11/29 Entry Date (DED): Update Date (DUPD): 1992/05/13

Field Availability:

Code Name		Occurrence
BRN	Beilstein Records	1
CN	Chemical Name	1
AUN	Autonomname	1
MF	Molecular Formula	1
FW	Formular Weight	1
LN	Lawson Number	. 3
CTYPE	Compound Type	1
CONSID	Constitution ID	1
TAUTID	Tautomer ID	1
BSO	Beilstein Citation	1
DED	Entry Date	1
DUPD	Update Date	1

MP Melting Point

1

This substance also occurs in Reaction Documents:

Code	Name	Occurrence
========		========
RX	Reaction Documents	. 1
RXPRO	Substance is Reaction Product	1

All References:

ALLREF

1. Patent: Shionogi and Co. Ltd. JP 7030335 1970, Chem. Abstr., 74(22872)

L17 ANSWER 3 OF 5 BEILSTEIN COPYRIGHT 2005 BEILSTEIN MDL on STN

Beilstein Records (BRN): 567960

Chemical Name (CN): (2,5-dimethyl-3-phenyl-pyrazolo<1,5-

a>pyrimidin-7-yl)-(2-piperidin-1-yl-ethyl)-

amine

Autonom Name (AUN): (2,5-dimethyl-3-phenyl-pyrazolo<1,5-

a>pyrimidin-7-yl)-(2-piperidin-1-yl-ethyl)-

amine

Molec. Formula (MF): C21 H27 N5 Molecular Weight (MW): 349.48

Lawson Number (LN): 30310, 24081, 3018

Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 542073
Tautomer ID (TAUTID): 560718
Beilstein Citation (BSO): 5-26

Entry Date (DED): 1988/11/28 Update Date (DUPD): 1992/09/03

Field Availability:

Code	Name Occur		
======			
BRN	Beilstein Records	1	
CN	Chemical Name	1	
AUN	Autonomname	1	
MF	Molecular Formula	1	
FW	Formular Weight	1	
LN	Lawson Number	3	
CTYPE	Compound Type	1	
CONSID	Constitution ID	. 1	
TAUTID	Tautomer ID	1	
BSO	Beilstein Citation	1	
DED	Entry Date	1	
DUPD	Update Date	1	
MP	Melting Point	1	

This substance also occurs in Reaction Documents:

Code	Name	Occurrence
========		
RX	Reaction Documents	1.
RXPRO	Substance is Reaction Product	1

All References:

ALLREF

1. Patent: Shionogi and Co.; Ltd. JP 7030335 1970, Chem. Abstr., 74(22872)

L17 ANSWER 4 OF 5 BEILSTEIN COPYRIGHT 2005 BEILSTEIN MDL on STN

Beilstein Records (BRN): Chemical Name (CN): N, N-dimethyl-glycine 2,5-dimethyl-3-phenylpyrazolo<1,5-a>pyrimidin-7-ylamide 2-dimethylamino-N-(2,5-dimethyl-3-phenyl-Autonom Name (AUN): pyrazolo<1,5-a>pyrimidin-7-yl)-acetamide C18 H21 N5 O Molec. Formula (MF): Molecular Weight (MW): 323.40 30310, 3379, 2817 Lawson Number (LN): Compound Type (CTYPE): heterocyclic Constitution ID (CONSID): 520141 Tautomer ID (TAUTID): 551098 Beilstein Citation (BSO): 5-26 Entry Date (DED): 1988/11/28 Update Date (DUPD): 1992/09/03

Field Availability:

Name	Occurrence	
Beilstein Records	1	
Chemical Name	· 1	
Autonomname	1	
Molecular Formula	1	
Formular Weight	1	
Lawson Number	3	
Compound Type	1	
Constitution ID	1	
Tautomer ID.	1	
Beilstein Citation	1	
Entry Date	1	
Update Date	1	
Melting Point	1	
	Beilstein Records Chemical Name Autonomname Molecular Formula Formular Weight Lawson Number Compound Type Constitution ID Tautomer ID. Beilstein Citation Entry Date Update Date	

This substance also occurs in Reaction Documents:

Code	Name	Occurrence
=======		========
RX	Reaction Documents	1
RXPRO	Substance is Reaction Product	1

All References:

ALLREF

1. Patent: Shionogi and Co.; Ltd. JP 7030335 1970, Chem.Abstr., 74(22872)

L17 ANSWER 5 OF 5 BEILSTEIN COPYRIGHT 2005 BEILSTEIN MDL on STN

Beilstein Records (BRN):

N'-(2,5-dimethyl-3-phenyl-pyrazolo<1,5-a>pyrimidin-7-yl)-N,N-dimethyl-ethane-1,2-Chemical Name (CN):

Autonom Name (AUN):

diamine
N-(2,5-dimethyl-3-phenyl-pyrazolo<1,5a>pyrimidin-7-yl)-N',N'-dimethyl-ethane-

Molec. Formula (MF):
Molecular Weight (MW):

1,2-diamine
C18 H23 N5
309.41

Lawson Number (LN): 30310, 3018, 2817
Compound Type (CTYPE): heterocyclic
Constitution ID (CONSID): 519770
Tautomer ID (TAUTID): 549248

Beilstein Citation (BSO): 5-26
Entry Date (DED): 1988/11/28
Update Date (DUPD): 1991/10/16

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Field Availability:

Code	e Name Oc	
=======		
BRN	Beilstein Records	1
CN	Chemical Name	1
AUN	Autonomname	1
MF	Molecular Formula	1
FW	Formular Weight	1
LN	Lawson Number	3
CTYPE	Compound Type	1
CONSID	Constitution ID	1
TAUTID	Tautomer ID	1
BSO	Beilstein Citation	1
DED	Entry Date	1
DUPD	Update Date	1
MP	Melting Point	1

This substance also occurs in Reaction Documents:

Code	Name		Occurrence
======			
RX	Reaction	Documents	1

RXPRO Substance is Reaction Product

1

All References:

ALLREF

1. Patent: Shionogi and Co.; Ltd. JP 7030335 1970, Chem. Abstr., 74(22872)

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STUCCTURE L8

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chain nodes :
   20 21
ring nodes :
   1 2 3 4 5 6 7 8 9 14 15 16 17 18 19
ring/chain nodes :
   10 11 12 13
chain bonds :
   2-21 4-10 8-20 9-17
ring/chain bonds :
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ring bonds :
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exact/norm bonds :
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Match level :
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20:CLASS 21:CLASS Element Count : Node 20: Limited C,C1-6 Node 21: Limited C,C1-6

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> d his full
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L4

L6

(FILE 'HOME' ENTERED AT 13:51:24 ON 24 AUG 2005)

FILE 'REGISTRY' ENTERED AT 13:51:42 ON 24 AUG 2005

STRUCTURE UPLOADED L122 SEA SSS SAM L1 L2

D SCA

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FILE 'CAPLUS' ENTERED AT 13:55:14 ON 24 AUG 2005

E US2002-083245/APPS E US2002-83245/APPS

1 SEA ABB=ON PLU=ON US2002-83245/AP SEL RN L4

FILE 'REGISTRY' ENTERED AT 13:56:12 ON 24 AUG 2005

D COST L5

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FILE 'STNGUIDE' ENTERED AT 14:04:07 ON 24 AUG 2005

FILE 'REGISTRY' ENTERED AT 14:09:13 ON 24 AUG 2005 435 SEA SSS FUL L1

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FILE 'REGISTRY' ENTERED AT 14:14:39 ON 24 AUG 2005

L8 STRUCTURE UPLOADED

L9 13 SEA SUB=L6 SSS SAM L8

D SCA

L10 212 SEA SUB=L6 SSS FUL L8
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SAVE TEMP L10 TRU245STRB/A

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L12

L13

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19 SEA ABB=ON PLU=ON L7 NOT L11

D SCA L4 TI

FILE 'MEDLINE, EMBASE, BIOSIS' ENTERED AT 14:20:49 ON 24 AUG 2005 L14 0 SEA ABB=ON PLU=ON L6

FILE 'STNGUIDE' ENTERED AT 14:23:46 ON 24 AUG 2005

FILE 'REGISTRY' ENTERED AT 14:24:50 ON 24 AUG 2005
L15 0 SEA ABB=ON PLU=ON L10 AND BEILSTEIN/LC NOT CAPLUS/LC

FILE 'BEILSTEIN' ENTERED AT 14:26:13 ON 24 AUG 2005

L16 6 SEA SSS FUL L8

L17 5 SEA ABB=ON PLU=ON L16 NOT RN/FA

FILE 'STNGUIDE' ENTERED AT 14:27:10 ON 24 AUG 2005

FILE 'REGISTRY' ENTERED AT 14:29:17 ON 24 AUG 2005

FILE 'CAPLUS' ENTERED AT 14:29:21 ON 24 AUG 2005 D STAT QUE L11 D IBIB ABS HITSTR L11 1-5

FILE 'REGISTRY' ENTERED AT 14:30:53 ON 24 AUG 2005

FILE 'CAPLUS' ENTERED AT 14:30:57 ON 24 AUG 2005

D STAT QUE L13

D IBIB ABS HITSTR L13 1-19

FILE 'REGISTRY' ENTERED AT 14:32:51 ON 24 AUG 2005

FILE 'MEDLINE' ENTERED AT 14:32:55 ON 24 AUG 2005

FILE 'EMBASE' ENTERED AT 14:33:00 ON 24 AUG 2005

FILE 'BIOSIS' ENTERED AT 14:33:04 ON 24 AUG 2005 D STAT QUE L14

FILE 'BEILSTEIN' ENTERED AT 14:33:37 ON 24 AUG 2005

D STAT QUE L17

D QRD ALLREF L17 1-5

FILE 'STNGUIDE' ENTERED AT 14:35:08 ON 24 AUG 2005

FILE HOME

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 23 AUG 2005 HIGHEST RN 861509-89-9 DICTIONARY FILE UPDATES: 23 AUG 2005 HIGHEST RN 861509-89-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See HELP SLIMITS for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

FILE CAPLUS

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FILE COVERS 1907 - 24 Aug 2005 VOL 143 ISS 9 FILE LAST UPDATED: 23 Aug 2005 (20050823/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE STNGUIDE
FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Aug 12, 2005 (20050812/UP).

FILE MEDLINE

FILE LAST UPDATED: 23 AUG 2005 (20050823/UP). FILE COVERS 1950 TO DATE.

On December 19, 2004, the 2005 MeSH terms were loaded.

The MEDLINE reload for 2005 is now available. For details enter HELP RLOAD at an arrow promt (=>). See also:

http://www.nlm.nih.gov/mesh/

http://www.nlm.nih.gov/pubs/techbull/nd04/nd04 mesh.html

OLDMEDLINE now back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2005 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE EMBASE

FILE COVERS 1974 TO 18 Aug 2005 (20050818/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BIOSIS

FILE COVERS 1969 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 17 August 2005 (20050817/ED)

FILE RELOADED: 19 October 2003.

FILE BEILSTEIN

FILE RELOADED ON OCTOBER 20, 2002

FILE LAST UPDATED ON JUNE 29, 2005

FILE COVERS 1771 TO 2005.

FILE CONTAINS 9,271,550 SUBSTANCES

>>>PLEASE NOTE: Reaction Data and substance data are stored in separate documents and can not be searched together in one query. Reaction data for BEILSTEIN compounds may be displayed immediately with the display codes PRE (preparations) and REA (reactions). A substance answer set retrieved after the search for a chemical name, a compounds with available reaction information by combining with PRE/FA, REA/FA or more generally with RX/FA. The BEILSTEIN Registry Number (BRN) is the link between a BEILSTEIN compound and belonging reactions. For mo detailed reaction searches BRNs can be searched as reaction partner BRNs Reactant BRN (RX.RBRN) or Product BRN (RX.PBRN).<<<

>>>	FOR	SEARCHING	PREPARATIONS	SEE	HELP	PRE	<<<
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- * PLEASE NOTE THAT THERE ARE NO FORMATS FREE OF COST.
- * SET NOTICE FEATURE: THE COST ESTIMATES CALCULATED FOR SET NOTICE
- * ARE BASED ON THE HIGHEST PRICE CATEGORY. THEREFORE; THESE
- * ESTIMATES MAY NOT REFLECT THE ACTUAL COSTS.
- * FOR PRICE INFORMATION SEE HELP COST

- * PATENT NUMBERS (PN) AND BABS ACCESSION NUMBERS (BABSAN) CAN NOW BE SEARCHED, SELECTED AND TRANSFERRED.
- * NEW DISPLAY FORMATS ALLREF, ALLP AND BABSAN SHOW ALL REFERENCES, ALL PATENT REFERENCES, OR ALL BABS ACCESSION NUMBERS FOR A COMPOUND AT A GLANCE.